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PTO

UTILITY PATENT APPLICATION TRANSMITTAL	
(Only for new nonprovisional applications under 37 CFR 1.53(b))	
Attorney Docket No	195976US0
First Inventor or Application Identifier	Reinhard SCHNEIDER, et al.
Title	LIGHTENING DYED TEXTILE MATERIAL

APPLICATION ELEMENTS See MPEP chapter 600 concerning utility patent application contents		ADDRESS TO: Assistant Commissioner for Patents Box Patent Application Washington, DC 20231	
<p>1. <input checked="" type="checkbox"/> Fee Transmittal Form (e.g. PTO/SB/17) (Submit an original and a duplicate for fee processing)</p> <p>2. <input checked="" type="checkbox"/> Specification Total Pages 18</p> <p>3. <input type="checkbox"/> Drawing(s) (35 U.S.C. 113) Total Sheets </p> <p>4. <input checked="" type="checkbox"/> Oath or Declaration Total Pages 3</p> <p>a. <input checked="" type="checkbox"/> Newly executed (original)</p> <p>b. <input type="checkbox"/> Copy from a prior application (37 C.F.R. §1.63(d)) (for continuation/divisional) with box 15 completed</p> <p>i. <input type="checkbox"/> DELETION OF INVENTOR(S) Signed statement attached deleting inventor(s) named in the prior application, see 37 C.F.R. §1.63(d)(2) and 1.33(b).</p> <p>5. <input type="checkbox"/> Incorporation By Reference (usable if box 4B is checked) The entire disclosure of the prior application, from which a copy of the oath or declaration is supplied under Box 4B, is considered to be part of the disclosure of the accompanying application and is hereby incorporated by reference therein.</p>		ACCOMPANYING APPLICATION PARTS <p>6. <input type="checkbox"/> Assignment Papers (cover sheet & document(s))</p> <p>7. <input type="checkbox"/> 37 C.F.R. §3.73(b) Statement <input type="checkbox"/> Power of Attorney (when there is an assignee)</p> <p>8. <input type="checkbox"/> English Translation Document (if applicable)</p> <p>9. <input checked="" type="checkbox"/> Information Disclosure Statement (IDS)/PTO-1449 <input checked="" type="checkbox"/> Copies of IDS Citations (5)</p> <p>10. <input type="checkbox"/> Preliminary Amendment</p> <p>11. <input checked="" type="checkbox"/> White Advance Serial No. Postcard</p> <p>12. <input type="checkbox"/> Small Entity Statement(s) <input type="checkbox"/> Statement filed in prior application. Status still proper and desired.</p> <p>13. <input type="checkbox"/> Certified Copy of Priority Document(s) (if foreign priority is claimed)</p> <p>14. <input checked="" type="checkbox"/> Other: <input type="checkbox"/> Statement of Relevancy, Notice of Priority</p>	
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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

INVENTOR(S) Reinhard SCHNEIDER, et al.

SERIAL NO: New Application

- FILING DATE: Herewith

FOR: LIGHTENING DYED TEXTILE MATERIAL

FEE TRANSMITTAL

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Lightening dyed textile material

This invention relates to a process for lightening or partially decolorizing vat- or sulfur-dyed textile materials, which comprises treating the material with 5 aminoalkanesulfinic acids in a neutral or weakly acidic medium, and to preparations of the aminoalkanesulfinic acids for carrying out this process.

10 In vat and sulfur dyeing, textile materials comprising cellulosic fibers, such as cotton, linen, hemp or regenerated cellulose fibers such as modal fibers or unmodified viscose rayon, with or without other, especially synthetic, fibers, eg. polyester fibers, for 15 example cotton textiles or cotton-polyester blend textiles, are treated with vatted vat or sulfur dyes in an alkaline medium. After the vatted dye has gone onto the cellulosic fiber, the textile material is rinsed neutral and then reoxidized.

20 Vatting is the conversion of certain water-insoluble dyes (vat dyes or sulfur dyes) by reduction in an alkaline medium into a water-soluble hydro or leuco compound whose anion has sufficient affinity for fiber composed of natural or regenerated cellulose, for 25 example cotton fiber or rayon. On reoxidation of the leuco compound, for example by exposure to air, the insoluble dye is re-formed in a very finely divided stage in the cellulosic fiber and thus ensures high-grade wash- and crock- and lightfastness.

30

Vat and sulfur dyes may be applied to textiles at every stage of processing, ie. not only to fabrics, such as

wovens, knits and nonwovens, but also to yarns. Textile materials are generally dyed in the form of the unprocessed fabrics; however, it is also perfectly possible to dye garments made of the textile materials 5 mentioned. Vat and sulfur dyes may also be applied by printing. In this case, thickened vats of vat or sulfur dyes are used and the printed material is subjected to a heat treatment, for example by steaming, and subsequently finished as in dyeing.

10

Yarn dyeing is predominantly practiced to produce textile sheet materials featuring special effects, for example bicolor effects. A well known textile material featuring such a special effect is denim. To produce 15 denim, the yarns which in the later woven fabric will form the warp threads which extend in the longitudinal direction of the woven web are dyed in a certain color, while the yarns for the transverse weft threads are dyed in a different color or are left undyed. The yarns 20 are then woven up on looms.

Hardwearing workwear items and, these days, blue jeans especially are generally manufactured using a coarse cotton fabric (denim), which is customarily dyed with 25 the vat dye indigo. To create the typical appearance of a pair of blue jeans, generally only the warp threads are dyed prior to weaving and the weft is left colorless. After weaving and making up, it is frequently desired that the garment be partially 30 decolorized again. In fact, only exposed areas of the garment are to be decolorized, the seams for example. Moreover, the pronounced color contrast between the dyed warp threads and the colorless weft threads is to be preserved, ie. the latter must not become stained in 35 the course of the lightening process. There is frequently also a desire to achieve a particular visual effect, for example the "used look" or that only

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exposed, comparatively highly stressed areas are lightened.

The partial subsequent removal of dye can be effected
5 for example mechanically by washing in the presence of pumice, enzymatically by the partial destruction of the cellulose, or chemically by altering or destroying the dye (stripping). A frequent choice is the stonewashing method, where the fabric is treated mechanically with
10 pumice and/or cellulases. Stonewashing is time-consuming and inconvenient, since the pumice stones have to be removed again from the material, and this is at least partly done by hand. The stones are responsible for abrading the equipment and major
15 amounts of sludge arise and have to be disposed of. The visual effect obtained in this way is very good, but the lightening effect is only minimal, so that generally a bleaching process is carried out in addition.

20 The bleach can be effected enzymatically, oxidatively or reductively.

An ecologically particularly advantageous bleach is the
25 enzymatic bleach by means of laccases which is described in WO 97 25 469. However, laccases are so costly that this process is too uneconomical for everyday textile finishing practice.¹

30 The oxidative removal of dyes may be carried out for example using strong oxidizing agents such as alkali metal hypochlorite, ozone or alkali metal permanganate. However, oxidative processes are disadvantageous because of the pronounced fiber damage and, especially
35 in the case of the use of hypochlorites, the unfavorable ecological aspects (AOX).

Despite these disadvantages, oxidative dye destruction today is customarily carried out using hypochlorite.

5 Reductive removal of vat and sulfur dyes may be effected using various reducing agents, albeit subject to specific restrictions if only partial removal of the dye is desired.

10 Very strong reducing agents (for example alkali metal dithionite or thiourea dioxide) are unsuitable, since they provide much too rapid and complete, in some instances instantaneous, vatting of many vat dyes, especially indigo, for example. As a result, much too much dye is stripped off in uncontrolled fashion. Weak
15 reducing agents, by contrast, such as glucose or hydroxyacetone, require a relatively high processing temperature, a high pH, ie. a relatively large amount of alkali metal (eg. sodium or potassium) hydroxide, and a high concentration of the reducing agent. In
20 addition, the treatment time is relatively long. Moreover, organic reducing agents, such as glucose, for example, are no longer acceptable in wastewater owing to their high chemical oxygen demand (COD); and the washing until neutral is very costly. Despite these
25 problems, relatively mildly reductive bleaching agents, for example, glucose, are still being used where necessary (JP 96-270034).

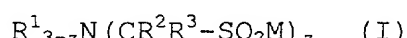
30 The vat dye detached from the warp should not stain the colorless weft threads, which tends to happen under reductive conditions, for example in the presence of alkaline glucose solution. It is therefore customary to include in the wash liquor a backstain inhibitor to control any backstaining by the dissolved dye because
35 the dissolved dye generally has greater affinity for the backstain inhibitor than for the fiber.

An improved reductive bleaching process, using hydroxyalkanesulfinic acids as bleaching agents, is described in DE-A-19 708 973. The disadvantage of this process is that it too has to be carried out in a 5 strongly alkaline medium with its attendant backstaining and the associated low color contrast.

It is an object of the present invention to provide a process for the controlled decolorizing or lightening 10 of vat- or sulfur-dyed textile materials whereby the aforementioned disadvantages are avoided or reduced and whereby an optimum color contrast (washdown effect) is achieved relatively consistently, rapidly and environmentally safely especially in the partial 15 decolorizing (lightening) of denim fabrics.

We have found that, surprisingly, this object is achieved by the textile finishing process of the 20 invention described hereinbelow.

The present invention accordingly provides a process 25 for controlled preferably partial, decolorization (lightening) of vat- or sulfur-dyed or -printed textile material, especially denim fabric, which comprises treating the textile material to be lightened or 30 decolorized with one or more compounds (aminoalkanesulfonates) of the formula I



35 where
z is 1, 2 or 3,
 R^1 is
a) when z is 1 or 2: hydrogen, alkyl of 1 to 18 carbon atoms or $HOCH_2CH_2$,
b) when z is 2: additionally OH, and

c) when z is 1: either as defined under a) independently for the two R^1 radicals or as defined under a) in one instance and as defined under b) in the other,

5 R^2 and R^3 , which may be the same or different, are each hydrogen or alkyl of 1 to 4 carbon atoms subject to the proviso that together they have not more than 4 carbon atoms, and

10 M is one equivalent of a mono- or divalent metal atom, at pH 4 - 7, followed if desired by an aftertreatment with hydrogen peroxide.

The process of the invention is preferably carried out using compounds of the formula I where z is 2 or 3 especially 3 and also compounds in which R^1 is hydrogen or alkyl R^1 has not more than 10 especially not more than 4 carbon atoms.

Preference is also given to compounds of formula I where R^2 and R^3 are independently hydrogen, methyl or ethyl and together have not more than 3 preferably only 2 carbon atoms. Preferred metals M are alkali and alkaline earth metals and zinc. Particular preference is given to compounds of this type in which a combination of the abovementioned preferred features is present, for example a compound of the formula I where R^1 , R^2 and R^3 are each hydrogen.

The compounds of the formula I can also be used as individuals. But it is more advantageous to use mixtures of these compounds in which the z indices have different meanings, especially mixtures in which the various compounds are present in that ratio which corresponds to their equilibrium concentration in an aqueous system of compounds of the formula I, the amine or the hydroxylamine of the formula $R^{13-z}NH_z$ and a hydroxyalkanesulfinate of the formula $HO-CR^2R^3-SO_2M$, where R^1 , R^2 , R^3 , z and M are each as defined above and

the molar ratio of sulfur-containing compounds to nitrogen-containing compounds is in the range from 0.2 to 1.1, preferably from 0.25 to 1.0, especially from 0.3 to 0.5.

5

It will be appreciated that the process of the invention can also be carried out using mixtures of compounds of the formula I which differ with regard to the meanings of R¹ and/or R² and/or R³.

10

The aminoalkanesulfonates of the formula I can also be used with advantage in mixture with the corresponding aminoalkanesulfonates, in which case these are present in an aminoalkanesulfinic acid:aminoalkanesulfonic acid ratio of from 3:1 to 1:3, preferably from 1.5:1 to 15 1:1.5 especially in a ratio of about 1:1.

In a further advantageous embodiment, the textile material is additionally and advantageously 20 concurrently treated with backstain inhibitors and/or dispersants and/or surfactants. These are used in total in an amount of from 0.5 to 10.0 g/l, preferably from 1 to 5 g/l. Backstain inhibitors are effective in particular in preventing the differently colored - or 25 in the case of denim the undyed - weft being (back)stained by the dissolved dye. Examples of backstain inhibitors are polyvinylpyrrolidone, naphthalenesulfonic acid-formaldehyde condensates, oleic acid alkoxylates and fatty acid alkoxylates.

30

The process of the invention is advantageously carried out at elevated temperature, preferably at 40 - 100°C, especially at 60 - 95°C, specifically at 75 - 90°C, under neutral or weakly acidic conditions, preferably 35 at a pH of from 4 to 7 especially from 5 to 7 and a liquor ratio of from 5:1 to 50:1 preferably from 10:1 to 20:1. The pH may be set using known acids, such as

citric acid, acetic acid or phosphoric acid. The pH may be stabilized by adding a buffer system eg. phosphoric acid/alkali metal phosphate or citric acid/alkali metal citrate, typically in an amount of about 2 g/l. The 5 compounds of the formula I are used in a concentration of from 0.001 to 0.3 mol/l, preferably from 0.01 to 0.1 mol/l, based on sulfinate.

Under the processing conditions described, the 10 bleaching time is generally in the range from 5 to 30 minutes, preferably in the range from 10 to 20 minutes, depending on the desired degree of lightening.

15 The stripping effect of the compounds of the formula I is simple to control. The degree of lightening increases with increasing temperature, with increasing bleaching agent concentration and with decreasing pH. Under strong bleaching conditions, the process of the 20 invention makes it possible to cut the stonewashing time appreciably.

Prior to the process of the invention being carried out, the material which is to be bleached is 25 advantageously desized. A customary enzymatic desizing process may be employed for this purpose, for example. Furthermore, the textile material may be subjected to a stonewashing process prior to bleaching. Stonewashing is customarily carried out using pumice and/or 30 cellulases. However, the bleach may also be carried out together with the stonewashing.

If desired, the textile material bleached according to the invention may additionally be subjected to an 35 oxidative aftertreatment. To this end, the bleached textile material may be treated for example at from 80 to 95°C and a liquor ratio of from 5:1 to 20:1 (eg.

10:1) with from 2 to 8 g/l (eg. 4 g/l) of 38°Bé sodium hydroxide, from 2 to 5 g/l of 50% by weight hydrogen peroxide and from 0.5 to 5 g/l (eg. 1 g/l) of backstain inhibitor, dispersant and/or surfactant for from 5 to 5 15 minutes (eg. 10 minutes).

In a further aftertreatment step, the textile material may be aftertreated as usual with softeners.

10 By way of further simplification of the application process, the solutions of the sulfonates or the solutions of the above-described equilibrium mixtures and/or their mixtures with the corresponding aminoalkanesulfonic acids may also be admixed with 15 process-specific assistants, for example the abovementioned backstain inhibitors, dispersants and/or surfactants. These preparations likewise form part of the subject-matter of the present invention.

20 The aminoalkanesulfonates of the formula I to be used for the bleaching process of the invention, the above-described equilibrium mixtures and the combination of these chemical entities with the corresponding aminoalkanesulfonates are also useful for preparing 25 discharge prints on dyeings of vat and sulfur dyes. For discharge printing, the neutral or weakly acidic solutions of the sulfonates or the solutions of the above-described equilibrium mixtures and/or their mixtures with the corresponding aminoalkanesulfonic acids are admixed with known thickeners and the thusly 30 obtained discharge print pastes are printed in the desired design on the dyeing to be discharged, subjected to a heat treatment and finished as usual. The process is particularly useful when discharge 35 printing is to be carried out in the neutral or weakly acidic pH range and/or when the desired discharge is not to white but half-tone patterns are to be produced.

Useful thickeners for preparing the discharge print pastes include all natural and synthetic substances known for preparing such print pastes, for example carob bean flour, salts of polyacrylic acid or solvent-based thickeners.

The aminoalkanesulfinic acid is prepared according to generally known processes (cf. EP-A-914516), for example by reacting sodium dithionite with 2 mol of an aldehyde or ketone of the general formula R^2COR^3 , where R^2 and R^3 are each as defined above, to obtain a mixture of hydroxyalkane-sulfinate and -sulfonate, from which the sulfinate may if desired be isolated for further reaction. The sulfinate obtained, but preferably the reaction mixture obtained from the reaction of the aldehyde or ketone with alkali metal dithionite, is condensed with ammonia, an amine or a hydroxylamine of the formula $R^1_{3-z}NH_z$, where R^1 and z are each as defined above, in a molar ratio of from 0.2 to 1.1, preferably from 0.25 to 1.0, especially from 0.3 to 0.5, in a suitable solvent, preferably in an aqueous medium, if desired at slightly elevated temperature. This produces a solution of the aminoalkanesulfinate or - if the hydroxyalkanesulfinate was not first isolated - of a mixture of aminoalkanesulfinate and aminoalkane-sulfonate, from which the aminoalkanesulfonate can be removed (cf. K. Reinking, E. Dehnel, H. Labhardt in Ber. 38, (1905), p. 1069-1080). Advantageously, however, the aminoalkanesulfonates are not isolated from the aqueous solution, nor the sulfonate separated off, and instead the as-obtained reaction mixture is used directly for the process of the invention. These solutions have very good stability in storage and the advantage for the user that the complicated dissolving of a solid substance in water is eliminated and instead the aqueous solution need only be diluted to use

concentration. In addition, automatic process control is made possible as a result.

The process of the invention is useful for the partial bleaching (lightening) of dyeings and prints with vat and sulfur dyes. Dyes whose dyeings or prints are bleachable according to the invention are listed in the Colour Index under vat dyes and sulfur dyes. Examples of what is bleachable by the process of the invention are dyeings with indigoid dyes, for example indigo, dibromoindigo, indigocarmine, tetrabromoindigo, tetrachloroindigo or thioindigo; with anthraquinonoid dyes, for example Indanthren Blue BC, Indanthren Brown NG, Indanthren Brilliant Green FFB, Indanthren Brilliant Orange GK, Indanthren Brilliant Orange GR, Indanthren Brilliant Orange RK, Indanthren Brilliant Rose R, Indanthren Brilliant Violet R extra, Indanthren Dark Blue BOA, Indanthren Golden Orange G, Indanthren Gray M, Indanthren Olive Green B, Indanthren Red RK, Indanthren Red FBB or Indanthren Reddish Violet RH; or with sulfur dyes, for example Immedial Pure Blue, with the Hydronblau products particularly important for blue workwear, Indocarbon CL and Sulfur Black T.

The treatment according to the process of the invention is particularly useful for dyeings with indigoid dyes, especially with indigo.

Compared with conventional processes, the bleaching process of the invention has a number of significant advantages.

The bleaching agents of the formula I have a reductive action and are therefore very gentle on the fiber. By working in a neutral to weakly acidic medium there is no need for the otherwise required costly and ecologically unfavorable neutralization.

The aminoalkanesulfinic acids of the formula I and their salts alone or in the embodiments more particularly described above are very useful for the 5 process of the invention, especially because their reactivity is between that of the weak reducing agents, such as glucose or hydroxyacetone, and that of the strong reducing agents, such as hydrosulfite (sodium dithionite) or thiourea dioxide. As a consequence, the 10 dyed textile material is generally decolorized to about 80 - 90%, especially under the conditions of the present invention, if desired nonuniformly.

15 The backstaining of dissolved dye, for example indigo, on the fibers especially on any differently colored - or in the case of denim material undyed - weft is minimal in that such a weft is left substantially unstained and, if appropriate, a very good contrast is obtained between warp dyeing and weft. The process of 20 the invention is hence a particularly efficient, simple and economical way of achieving the washdown effect. This makes the process of the invention especially useful in jeans washing.

25 Another surprise is that the portion of the vat or sulfur dye, especially the portion of indigo, which has become redeposited on the fiber is easily removable (for example, by an oxidative aftertreatment with hydrogen peroxide) when the bleach is carried out 30 according to the invention. This removal of indigo deposited on the fiber takes place even though hydrogen peroxide alone is not capable of lightening indigo dyeings.

35 The oxidative aftertreatment with hydrogen peroxide is thus a preferred embodiment of the present invention, which is preferably employed when backstaining is to be

minimal, when a concentrated liquor is to be used or when a very high degree of lightening is to be achieved in a single wash.

5 The invention further provides for the use of aminoalkanesulfinates of the formula I or of the above-described mixtures thereof for, preferably partial, decolorization (lightening) of vat- or sulfur-dyed textile materials.

10

The present invention further provides for the use of aminoalkanesulfinates of the formula I or of the above-described mixtures thereof for preparing discharge prints especially in the half-tone area.

15

The example which follows illustrates the invention.

Example

20 A drum washer is charged with 5 kg of desized and stonewashed jean pants, 100 l of water, 2 g/l of an oleic acid ethoxylate and 1.5 g/l of glacial acetic acid, started, heated to 85°C and at 85°C with 15 ml of a 50% by weight solution containing equimolar amounts
25 of hydroxymethanesulfinate condensed with 0.33 mol% of ammonia and hydroxymethanesulfonate condensed with 0.33 mol% of ammonia. The pH is 6.2 and in the course of the bleaching process it rises to 6.5¹ over 15 minutes. Thereafter the bleaching liquor is dropped hot from the
30 rotating drum and the textile material is rinsed once with cold water. Some specimens of the pants thus treated are dried, while the others are treated in a liquor ratio of 10:1 with a liquor containing 5 g/l of 50% by weight hydrogen peroxide, 1 g/l of oleic acid ethoxylate and 4 ml of 38°Bé sodium hydroxide solution
35 at 90°C for 10 minutes. This is followed by a single rinse with water containing 0.5 g/l of citric acid and

3 g/l of a commercially available soft hand agent, and drying.

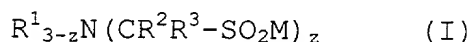
All the jean pants treated in the manner described
5 exhibit a conspicuous used look and very good contrast
between weft and warp threads. There is no sign
whatsoever of damage to the fabric. The inside pockets
of the pants not given the oxidative aftertreatment are
10 slightly blue, while the inside pockets of the pants
aftertreated with hydrogen peroxide are perfectly
white.

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August 10, 2000
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We claim:-

5 1. A process for controlled partial decolorization of
vat- or sulfur-dyed or -printed textile material,
especially denim fabric, which comprises treating
the textile material to be lightened or de-
colorized with one or more compounds (aminoalkane-
10 sulfonates) of the formula I



Where

15 z is 1, 2 or 3,

R^1 is

a) when z is 1 or 2: hydrogen, alkyl of 1 to 18
carbon atoms or $HOCH_2CH_2$,

b) when z is 2: additionally OH, and

20 c) when z is 1: either as defined under a)
independently for the two R^1 radicals
or as defined under a) in one instance and as
defined under b) in the other,

25 R^2 and R^3 , which may be the same or different, are
each hydrogen or alkyl of 1 to 4 carbon atoms
subject to the proviso that together they have not
more than 4 carbon atoms, and

M is one equivalent of a mono- or divalent metal
atom,

30 at pH 4 - 7, followed if desired by an
aftertreatment with hydrogen peroxide.

2. A process as claimed in claim 1, wherein in one or
more compounds of the formula I used z is 3 and R^1 ,
35 R^2 and R^3 are each hydrogen.

3. A process as claimed in claim 1, utilizing mixtures of compounds of the formula I where the z indices have different meanings, especially
5 mixtures in which the various compounds are present in that ratio which corresponds to their equilibrium concentration in an aqueous system of compounds of the formula I, the amine or the hydroxylamine of the formula $R^{1,3-z}NH_2$ and a hydroxyalkanesulfinate of the formula $HO-CR^2R^3-SO_2M$, where R1, R2, R3, z and M are each as defined above and the molar ratio of sulfur-containing compounds to nitrogen-containing compounds is in
10 the range from 0.2 to 1.1.

15

4. A process as claimed in claim 1, utilizing mixtures of compounds of the formula I with the corresponding aminoalkanesulfonates where the ratio of aminoalkanesulfinic acid to aminoalkanesulfonic acid is from about 3:1 to about 1:3.
20

25

5. A process as claimed in claim 1, wherein the textile material is additionally, preferably concurrently, treated with one or more further assistants from the group of the backstain inhibitors and/or dispersants and/or surfactants, preferably in total in an amount of from 0.5 to 10.0 g/l.

6. A process as claimed in claim 5, wherein the ratio of an aminoalkanesulfonate to further assistants is in the range from 20:1 to 5:1.
30

35

7. A process as claimed in claim 5, wherein the backstain inhibitor is polyvinylpyrrolidone, oleic acid alkoxylate or fatty acid alkoxylate.

8. A process as claimed in claim 1, wherein the compounds of the formula I are printed in the form of a print paste onto the dyeing to be bleached.

5 9. A preparation for carrying out the process of claim 1, comprising a solution of the sulfinites of the formula I and/or an equilibrium mixture as set forth in claim 3 and/or a mixture with the corresponding aminoalkanesulfonic acids as set forth in claim 4, and also process-specific assistants, especially backstain inhibitors, dispersants and/or surfactants.

10. 15. A method of using aminoalkanesulfinites of the formula I or of mixtures thereof, where the z indices have different meanings, for, preferably partial, decolorization (lightening) of vat- or sulfur-dyed textile materials.

20. 11. A method of using aminoalkanesulfinites of the formula I or of mixtures thereof, where the z indices have different meanings, for preparing discharge prints especially in the half-tone area.

Abstract

Lightening dyed textile material

5 Described is a process for lightening or partially decolorizing vat- or sulfur-dyed textile materials, which comprises treating the material with aminoalkanesulfinic acids in a neutral or weakly acidic medium, and preparations of the aminoalkanesulfinic
10 acids for carrying out this process.

Declaration, Power of Attorney

Page 1 of 3

0050/050599

We (I), the undersigned inventor(s), hereby declare(s) that:

My residence, post office address and citizenship are as stated below next to my name,

We (I) believe that we are (I am) the original, first, and joint (sole) inventor(s) of the subject matter which is claimed and for which a patent is sought on the invention entitled

Lightening dyed textile material

the specification of which

is attached hereto.

was filed on _____ as

Application Serial No. _____

and amended on _____.

was filed as PCT international application

Number _____

on _____

and was amended under PCT Article 19

on _____ (if applicable).

We (I) hereby state that we (I) have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

We (I) acknowledge the duty to disclose information known to be material to the patentability of this application as defined in Section 1.56 of Title 37 Code of Federal Regulations.

We (I) hereby claim foreign priority benefits under 35 U.S.C. § 119(a)-(d) or § 365(b) of any foreign application(s) for patent or inventor's certificate, or § 365(a) of any PCT International application which designated at least one country other than the United States, listed below and have also identified below, by checking the box, any foreign application for patent or inventor's certificate, or PCT International application having a filing date before that of the application on which priority is claimed. Prior Foreign Application(s)

Application No.	Country	Day/Month/Year	Priority Claimed
19940068.7	Germany	24 August 1999	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No

We (I) hereby claim the benefit under Title 35, United States Codes, § 119(e) of any United States provisional application(s) listed below.

(Application Number)	(Filing Date)
(Application Number)	(Filing Date)

We (I) hereby claim the benefit under 35 U.S.C. § 120 of any United States application(s), or § 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

Application Serial No.	Filing Date	Status (pending, patented, abandoned)

And we (I) hereby appoint: Norman F. Oblon, Registration Number 24, 618; Marvin J. Spivak, Registration Number 24, 913; Gregory J. Maier, Registration Number 25, 599; William E. Beaumont, Registration Number 30, 996; Steven B. Kelber, Registration Number 30, 073; Jean-Paul Lavallee, Registration Number 31, 451; Timothy R. Schwartz, Registration Number 32, 171; Stephen G. Baxter, Registration Number 32, 884; Richard L. Treanor, Registration Number 36, 379; Robert W. Hahl, Registration Number 33, 893, our (my) attorneys, with full powers

of substitution and revocation, to prosecute this application and to transact all business in the Patent Office connected therewith; and we (I) hereby request that all correspondence regarding this application be sent to the firm of **OBLON, SPIVAK, McCLELLAND, MAIER & NEUSTADT, P. C.**, whose Post Office Address is: Fourth Floor, 1755 Jefferson Davis Highway, Arlington, Virginia 22202.

We (I) declare that all statements made herein of our (my) own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Reinhard Schneider
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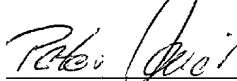


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